## **AMENDMENTS TO THE CLAIMS**

This listing of claims will replace all prior versions, and listings, of claims in the application:

Claim 1 (original): A method of treating a blood product which contains a nucleic acidcontaining pathogen to be inactivated, said method comprising

- a) adding psoralen to the blood product;
- b) irradiating the psoralen and the blood product to form a mixture comprising said blood product, free psoralen, and low molecular weight psoralen photoproducts; and
- c) contacting said mixture with a hypercrosslinked resin to remove at least substantially all of said free psoralen and said low molecular weight psoralen photoproducts.

Claim 2 (currently amended): The method of claim  $\theta$  1 wherein said psoralen comprises an aminopsoralen selected from the group consisting of 4'-primary amino-substituted psoralen and 5'-primary amino-substituted psoralen.

Claim 3 (currently amended): The method of claim  $\theta$  1 wherein said blood product comprises plasma.

Claim 4 (currently amended): The method of claim  $\theta$  1 wherein said hypercrosslinked resin is not pre-wetted prior to said act of contacting said mixture with said hypercrosslinked resin.

Claim 5 (currently amended): The method of claim  $\theta$  1 wherein said hypercrosslinked resin comprises a polyaromatic resin that is capable of adsorbing said free psoralen and said low molecular weight psoralen photoproducts.

Claim 6 (original): The method of claim 5 wherein said psoralen comprises an aminopsoralen selected from the group consisting of 4'-primary amino-substituted psoralen and 5'-primary amino-substituted psoralen.

Claim 7 (original): The method of claim 6 wherein said aminopsoralen comprises 4'-(4-amino-2-oxa)butyl-4,5',8-trimethylpsoralen.

Claim 8 (original): A method of removing free psoralen from a biological fluid comprising blood or a blood product, said free psoralen having been exposed to light having a wavelength that causes psoralen to covalently bind to a nucleic acid, the method comprising contacting said biological fluid with a hypercrosslinked adsorbent resin that is capable of removing said free psoralen; and removing at least substantially all of said free psoralen from said biological fluid with said hypercrosslinked adsorbent resin.

Claim 9 (original): The method of claim 8 wherein said resin is selected from the group consisting of: a polyaromatic resin having a mean surface area of about  $1100 \text{ m}^2/\text{gm}$ , a mean pore diameter of about 46Å, and a mesh size of about  $20\text{-}50\mu\text{m}$ ; a polyaromatic resin having a mean surface area of about  $725 \text{ m}^2/\text{gm}$ , a mean pore diameter of about 40Å, and a mesh size of about  $20\text{-}60\mu\text{m}$ ; and a functionalized polyaromatic resin having a mean surface area of about  $800 \text{ m}^2/\text{gm}$ , a mean pore diameter of about 25Å, and a mesh size of about  $20\text{-}50\mu\text{m}$ .

Claim 10 (original): The method of claim 8 wherein said biological fluid comprises a plasma blood product.

Claim 11 (original): The method of claim 8 wherein said biological fluid comprises a platelet-containing blood product.

Claim 12 (original): The method of claim 11 wherein said biological fluid further comprises a synthetic medium containing phosphate.

Claim 13 (original): The method of claim 8 wherein said resin is not pre-wetted prior to contacting said biological fluid with said resin.

Claim 14 (original): The method of claim 8 wherein said psoralen comprises an aminopsoralen selected from the group consisting of 4'-primary amino-substituted psoralen and 5'-primary amino-substituted psoralen.

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Claim 15 (original): The method of claim 14 wherein said aminopsoralen comprises 4'-(4-amino-2-oxa)butyl-4,5',8-trimethylpsoralen.

Claim 16 (original): The method of claim 8 wherein said hypercrosslinked resin comprises a hypercrosslinked polyaromatic resin.

Claim 17 (original): The method of claim 16 wherein said biological fluid is selected from the group consisting of plasma and platelets.

Claim 18 (original): The method of claim 16 wherein said psoralen comprises an aminopsoralen selected from the group consisting of 4'-primary amino-substituted psoralen and 5'-primary amino-substituted psoralen.

Claim 19 (original): The method of claim 16 wherein said psoralen comprises a brominated psoralen.

Claim 20 (original): The method of claim 16 wherein the biological fluid further comprises psoralen photo products, and wherein said resin additionally removes at least substantially all of said psoralen photo products.

Claim 21 (currently amended): A biological fluid formed by the method of claim  $\theta \underline{1}$ .

Claim 22 (original): A biological fluid formed by the method of claim 3.

Claim 23 (original): A biological fluid formed by the method of claim 8.

Claim 24 (original): A biological fluid formed by the method of claim 12.